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RESPIRATORY PATHOLOGY
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Submitted herewith is the Verified English Translation of French patent application no. 96 09 403, filed on July 19, 1996, to which the above-captioned application claims priority.

No fee is believed to be due for entry and consideration of this paper, however, any required fee may be charged to Deposit Account No. 50-0320.

Respectfully submitted,

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#9

IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

In the Application of : Audonnet et al.)
Serial No. 10/085,519)
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POLYNUCLEOTIDE VACCINE FORMULA IN PARTICULAR AGAINST
BOVINE RESPIRATORY PATHOLOGY

5 The present invention relates to a vaccine formula allowing the vaccination of bovines in particular against respiratory pathology. It also relates to a corresponding method of vaccination.

10 All bovines are carriers of viruses and bacteria which are potentially pathogenic in widely variable degrees.

15 Viruses can multiply when the specific immunity is weakened and when there are lesions of the respiratory tract. They are then excreted by the animal and may then contaminate other animals.

20 Among the viruses which are encountered, there may be mentioned in particular the type 3 parainfluenza virus (PI-3), of moderate inherent pathogenicity, the bovine respiratory syncytial virus (RSV) and the bovine herpesvirus (BHV) also called infectious bovine rhinotracheitis (IBR) virus, of high inherent pathogenicities.

25 Another virus which is particularly important for its immunodepressant role and its harmful effects on reproduction is the mucosal disease virus or bovine pestivirus (BVDV).

30 These viruses generally manifest themselves by a primary phase of hyperthermia, flu syndrome and respiratory disorders, with digestive disorders (diarrhoeas) in the case of BVD. This phase may be accompanied by a secondary phase with the onset of bronchopneumonia linked to bacterial, in particular *Pasteurella*, infections which can lead to death. This phenomenon is exacerbated in particular by the immunodepression resulting from BVD infection or by the 35 infection of macrophages by PI-3. Other symptoms may further appear, such as abortions with BVD and BHV.

It therefore appears necessary to try to develop an effective prevention against the principal

viruses involved in bovine respiratory pathology.

Associations of vaccines against certain viruses responsible for bovine respiratory pathology have already been proposed in the past.

5 The associations developed so far were prepared from inactivated vaccines or live vaccines and, optionally, mixtures of such vaccines. Their development poses problems of compatibility between valencies and of stability. It is indeed necessary to ensure both the
10 compatibility between the different vaccine valencies, whether from the point of view of the different antigens used or from the point of view of the formulations themselves, especially in the case where both inactivated vaccines and live vaccines are combined. The
15 problem of the conservation of such combined vaccines and also of their safety especially in the presence of an adjuvant also exists. These vaccines are in general quite expensive.

20 Patent Applications WO-A-90 11092, WO-A-93 19183, WO-A-94 21797 and WO-A-95 20660 have made use of the recently developed technique of polynucleotide vaccines. It is known that these vaccines use a plasmid capable of expressing, in the host's cells, the antigen inserted into the plasmid. All the routes of
25 administration have been proposed (intraperitoneal, intravenous, intramuscular, transcutaneous, intra-dermal, mucosal and the like). Various vaccination means can also be used, such as DNA deposited at the surface of gold particles and projected so as to penetrate into the animal's skin (Tang et al., Nature 356, 152-154, 1992) and liquid jet injectors which make it possible to transfect at the same time the skin, the muscle, the fatty tissues and the mammary tissues
30 (Furth et al., Analytical Biochemistry, 205, 365-368, 1992).
35

The polynucleotide vaccines may also use both naked DNAs and DNAs formulated, for example, inside cationic lipid or liposomes.

G.J.M. COX has already proposed polynucleotide

vaccination against type 1 bovine herpes virus in J. of Virology, Volume 67, No. 9, September 1993, 5664-5667. The authors have also described plasmids integrating the gI (gB), gIII (gC) and gIV (gD) genes.

5 In Vaccine, Volume 13, No. 4, 415-421, 1995, J.E. CROWE presents a general review of the different methods of vaccination against respiratory syncytial virus and against type 3 parainfluenza virus. This review reexamines all the possibilities offered by the
10 current vaccination techniques and simply suggests that the polynucleotide immunization technique could be useful in the immunization strategy against RSV and PI-3. No plasmid construction or result of vaccination of bovines against these viruses is described in this
15 document.

20 The invention therefore proposes to provide a multivalent vaccine formula which makes it possible to ensure vaccination against a number of pathogenic viruses involved in particular in bovine respiratory pathology and thus to ensure effective vaccination against this pathology.

25 Another objective of the invention is to provide such a vaccine formula combining different valencies while exhibiting all the criteria required for mutual compatibility and stability of the valencies.

30 Another objective of the invention is to provide such a vaccine formula which makes it possible to combine different valencies in the same vehicle.

35 Another objective of the invention is to provide such a vaccine formula which is easy and inexpensive to use.

 Yet another objective of the invention is to provide such a vaccine formula and a method for vaccinating bovines which make it possible to obtain a multivalent protection with a high level of efficiency and of long duration, as well as good safety and an absence of residues.

 The subject of the present invention is there-

fore a vaccine formula in particular against bovine respiratory pathology, comprising at least three polynucleotide vaccine valencies each comprising a plasmid integrating, so as to express it in vivo in the host cells, a gene with one bovine respiratory pathogen valency, these valencies being selected from the group consisting of bovine herpesvirus, bovine respiratory syncytial virus, mucosal disease virus and type 3 parainfluenza virus, the plasmids comprising, for each valency, one or more of the genes selected from the group consisting of gB and gD for the bovine herpesvirus, F and G for the bovine respiratory syncytial virus, E2, C + E1 + E2 and E1 + E2 for the mucosal disease virus, and HN and F for the type 3 parainfluenza virus.

Valency in the present invention is understood to mean at least one antigen providing protection against the virus for the pathogen considered, it being possible for the valency to contain, as subvalency, one or more modified or natural genes from one or more strains of the pathogen considered.

Pathogenic agent gene is understood to mean not only the complete gene but also the various nucleotide sequences, including fragments which retain the capacity to induce a protective response. The notion of a gene covers the nucleotide sequences equivalent to those described precisely in the examples, that is to say the sequences which are different but which encode the same protein. It also covers the nucleotide sequences of other strains of the pathogen considered, which provide cross-protection or a protection specific for a strain or for a strain group. It also covers the nucleotide sequences which have been modified in order to facilitate the in vivo expression by the host animal but encoding the same protein.

Preferably, the vaccine formula according to the invention comprises the four valences.

As regards the BHV valency, use is preferably made of the two genes encoding gB and gD, in different

plasmids or in one and the same plasmid. Optionally, but less preferably, either of these genes can be used.

For the RSV valency, use is preferably made of the two G and F genes integrated into two different 5 plasmids or into one and the same plasmid. Optionally, but less preferably, the F gene can be used alone.

For the BVD valency, use will preferably be made of a plasmid integrating the E2 gene. Optionally, but less preferably, a plasmid coding for E1 and E2 10 together or for the combination consisting of C, E1 and E2 can be used.

For the PI-3 valency, use is preferably made of the combination of the two HN and F genes in two different plasmids or in one and the same plasmid. It 15 is also possible to use only the HN gene.

A preferred vaccine formula according to the invention comprises and ensures the expression of the BHV gB and gD genes, the RSV G and F genes, the BVD E2 genes and PI-3 HN and F genes.

20 The vaccine formula according to the invention can be provided in a dose volume of between 0.1 and 10 ml and in particular between 1 and 5 ml.

The dose will be generally between 10 ng and 1 mg, preferably between 100 ng and 500 µg and more preferably 25 between 1 µg and 250 µg per plasmid type.

Use will preferably be made of naked plasmids simply placed in the vaccination vehicle which will be in general saline (0.9% NaCl), ultrapure water, TE buffer and the like. All the polynucleotide 30 vaccine forms described in the prior art can of course be used.

Each plasmid comprises a promoter capable of ensuring the expression of the gene inserted, under its control, into the host cells. This will be in general a 35 strong eukaryotic promoter and in particular a cytomegalovirus early CMV-IE promoter of human or murine origin, or optionally of another origin such as rats, pigs and guinea pigs.

More generally, the promoter may be either of

5 viral origin or of cellular origin. As viral promoter other than CMV-IE, there may be mentioned the SV40 virus early or late promoter or the Rous sarcoma virus LTR promoter. It may also be a promoter from the virus from which the gene is derived, for example the gene's own promoter.

10 As cellular promoter, there may be mentioned the promoter of a cytoskeleton gene, such as for example the desmin promoter (Bolmont et al., Journal of Submicroscopic Cytology and Pathology, 1990, 22, 117-122; and Zhenlin et al., Gene, 1989, 78, 243-254), or alternatively the actin promoter.

15 When several genes are present in the same plasmid, these may be presented in the same transcription unit or in two different units.

20 The combination of the different vaccine valencies according to the invention may be preferably achieved by mixing the polynucleotide plasmids expressing the antigen(s) of each valency, but it is also possible to envisage causing antigens of several valencies to be expressed by the same plasmid.

25 The subject of the invention is also monovalent vaccine formulae comprising one or more plasmids encoding one or more genes from one of the viruses selected from the group consisting of BRSV, BVD and PI-3, the genes being those described above. Besides their monovalent character, these formulae may possess the characteristics stated above as regards the choice of the genes, their combinations, the composition of the plasmids, the dose volumes, the doses and the like.

30 The monovalent vaccine formulae may be used (i) for the preparation of a polyvalent vaccine formula as described above, (ii) individually against the actual pathology, (iii) combined with a vaccine of another type (live or inactivated whole, recombinant, subunit) against another pathology, or (iv) as booster for a vaccine as described below.

35 The subject of the present invention is in fact also the use of one or more plasmids according to the

invention for the manufacture of a vaccine intended to vaccinate bovines first vaccinated by means of a first conventional vaccine of the type in the prior art, in particular, selected from the group consisting of a live 5 whole vaccine, an inactivated whole vaccine, a subunit vaccine, a recombinant vaccine, this first vaccine having, that is to say containing or capable of expressing, the antigen(s) encoded by the plasmids or antigen(s) providing cross-protection.

10 Remarkably, the polynucleotide vaccine has a potent booster effect which results in an amplification of the immune response and the acquisition of a long-lasting immunity.

15 In general, the first-vaccination vaccines can be selected from commercial vaccines available from various veterinary vaccine producers.

20 The subject of the invention is also a vaccination kit grouping together a first-vaccination vaccine as described above and a vaccine formula according to the invention for the booster. It also relates to a vaccine formula according to the invention accompanied by a leaflet indicating the use of this formula as a booster for a first vaccination as described above.

25 The subject of the present invention is also a method for vaccinating bovines against respiratory pathology, comprising the administration of the effective vaccine formula as described above. This vaccination method comprises the administration of one 30 or more doses of the vaccine formula, it being possible for these doses to be administered in succession over a short period of time and/or in succession at widely spaced intervals.

35 The vaccine formulae according to the invention can be administered, in the context of this method of vaccination, by the different routes of administration proposed in the prior art for polynucleotide vaccination and by means of known techniques of administration.

The subject of the invention is also the method

of vaccination consisting in making a first vaccination as described above and a booster with a vaccine formula according to the invention.

5 In a preferred embodiment of the process according to the invention, there is administered in a first instance, to the animal, an effective dose of the conventional type vaccine especially inactivated, live, attenuated or recombinant, type, or alternatively a subunit vaccine, so as to provide a first vaccination, 10 and, within a period preferably of 2 to 6 weeks, the polyvalent or monovalent vaccine according to the invention is administered.

15 The invention also relates to the method of preparing the vaccine formulae, namely the preparation of the valencies and mixtures thereof, as evident from this description.

The invention will now be described in greater detail with the aid of the embodiments of the invention taken with reference to the accompanying drawings.

20

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30 Figure No. 9 : Plasmid pAB060
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Sequence listing SEQ ID No.

35 SEQ ID No. 1 : Sequence of the BHV-1 gB gene
(strain ST)
SEQ ID No. 2 : Oligonucleotide PB234
SEQ ID No. 3 : Oligonucleotide PB235
SEQ ID No. 4 : Oligonucleotide AB162

SEQ ID No. 5 : Oligonucleotide AB163
SEQ ID No. 6 : Oligonucleotide AB026
SEQ ID No. 7 : Oligonucleotide AB027
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5 SEQ ID No. 9 : Oligonucleotide AB029
SEQ ID No. 10 : Oligonucleotide AB110
SEQ ID No. 11 : Oligonucleotide AB111
SEQ ID No. 12 : Oligonucleotide AB114
SEQ ID No. 13 : Oligonucleotide AB115
10 SEQ ID No. 14 : Oligonucleotide AB116
SEQ ID No. 15 : Oligonucleotide AB117
SEQ ID No. 16 : Oligonucleotide AB130
SEQ ID No. 17 : Oligonucleotide AB131
SEQ ID No. 18 : Oligonucleotide AB132
15 SEQ ID No. 19 : Oligonucleotide AB133

EXAMPLES

Example 1: Culture of the viruses

20 The viruses are cultured on the appropriate cellular system until a cytopathic effect is obtained. The cellular systems to be used for each virus are well known to persons skilled in the art. Briefly, the cells sensitive to the virus used, which are cultured in 25 Eagle's minimum essential medium (MEM medium) or another appropriate medium, are inoculated with the viral strain studied using a multiplicity of infection of 1. The infected cells are then incubated at 37°C for the time necessary for the appearance of a complete cytopathic 30 effect (on average 36 hours).

Example 2: Extraction of the viral genomic DNAs

After culturing, the supernatant and the lysed cells are harvested and the entire viral suspension is 35 centrifuged at 1000 g for 10 minutes at +4°C so as to remove the cellular debris. The viral particles are then harvested by ultracentrifugation at 400,000 g for 1 hour at +4°C. The pellet is taken up in a minimum volume of buffer (10 mM Tris, 1 mM EDTA). This concentrated viral

suspension is treated with proteinase K (100 µg/ml final) in the presence of sodium dodecyl sulphate (SDS) (0.5% final) for 2 hours at 37°C. The viral DNA is then extracted with a phenol/chloroform mixture and then precipitated with 2 volumes of absolute ethanol. After leaving overnight at -20°C, the DNA is centrifuged at 10,000 g for 15 minutes at +4°C. The DNA pellet is dried and then taken up in a minimum volume of sterile ultrapure water. It can then be digested with restriction enzymes.

Example 3: Isolation of the viral genomic RNAs

The RNA viruses were purified according to techniques well known to persons skilled in the art. The genomic viral RNA of each virus was then isolated using the "guanidium thiocyanate/phenol-chloroform" extraction technique described by P. Chomczynski and N. Sacchi (Anal. Biochem., 1987, 162, 156-159).

Example 4: Molecular biology techniques

All the constructions of plasmids were carried out using the standard molecular biology techniques described by J. Sambrook et al. (*Molecular Cloning: A Laboratory Manual*, 2nd Edition, Cold Spring Harbor Laboratory, Cold Spring Harbor, New York, 1989). All the restriction fragments used for the present invention were isolated using the "Geneclean" kit (BIO 101 Inc. La Jolla, CA).

Example 5: RT-PCR technique

Specific oligonucleotides (comprising restriction sites at their 5' ends to facilitate the cloning of the amplified fragments) were synthesized such that they completely cover the coding regions of the genes which are to be amplified (see specific examples). The reverse transcription (RT) reaction and the polymerase chain reaction (PCR) were carried out according to standard techniques (J. Sambrook et al., *Molecular Cloning: A Laboratory Manual*, 2nd Edition, Cold Spring Harbor

Laboratory, Cold Spring Harbor, New York, 1989). Each RT-PCR reaction was performed with a pair of specific amplimers and taking, as template, the viral genomic RNA extracted. The complementary DNA amplified was extracted 5 with phenol/chloroform/isoamyl alcohol (25:24:1) before being digested with restriction enzymes.

Example 6: plasmid pVR1012

10 The plasmid pVR1012 (Figure No. 1) was obtained from Vical Inc., San Diego, CA, USA. Its construction has been described in J. Hartikka et al. (Human Gene Therapy, 1996, 7, 1205-1217).

15 **Example 7: Construction of the plasmid pPB156 (BHV-1 gB gene)**

The BHV-1 bovine herpesvirus (ST strain) genomic DNA (Leung-Tack P. et al. Virology, 1994, 199, 409-421) was prepared according to the technique described in Example 2 and was digested with BamHI. 20 After purification, the 18 kbp BamHI-BamHI fragment was cloned into the vector pBR322, previously digested with BamH1, to give the plasmid pIBR-4-BamHI (22 kbp).

25 The plasmid pIBR-4-BamHI was then digested with SalI in order to liberate a 6.6 kbp SalI-SalI fragment containing the gene encoding the BHV-1 gB glycoprotein (Figure No. 2 and SEQ ID No. 1). This fragment was cloned into the vector pBR322, previously digested with SalI, to give the plasmid pIBR-6,6-SalI (10.9 kbp).

30 The plasmid pIBR-6,6-SalI was digested with NheI and BglII in order to liberate a 2676 bp NheI-BglII fragment containing the gene encoding the bovine herpesvirus (BHV-1) gB glycoprotein (fragment A).

35 A PCR reaction was carried out with the genomic DNA from the bovine herpesvirus (BHV-1) (ST strain) and with the following oligonucleotides:

PB234 (30 mer) (SEQ ID No. 2)

5' TTGTCGACATGGCCGCTCGCGGCGGTGCTG 3'

PB235 (21 mer) (SEQ ID No. 3)

5' GCAGGGCAGCGGCTAGCGCGG 3'

so as to isolate the 5' part of the gene encoding the BHV-1 gB glycoprotein. After purification, the 153 bp PCR product was digested with SalI and NheI in order to isolate a 145 bp SalI-NheI fragment (fragment B).

5 The fragments A and B were ligated together with the vector pVR1012 (Example 6), previously digested with SalI and BamHI, to give the plasmid pPB156 (7691 bp) (Figure No. 3).

10 **Example 8: Construction of the plasmid pAB087 (BHV-1 gD gene)**

A PCR reaction was carried out with the genomic DNA from the bovine herpesvirus (BHV-1) (ST strain) (P. Leung-Tack et al., Virology, 1994, 199, 409-421), prepared according to the technique described in Example 2, and with the following oligonucleotides:

15 AB162 (31 mer) (SEQ ID No. 4)

5' AAACTGCAGATGCAAGGGCCGACATTGGCCG 3'

20 AB163 (30 mer) (SEQ ID No. 5)

5' ATCTTGTACCATATGACCGTGGCGTTG 3'

so as to amplify the 5' part of the gene encoding the bovine herpesvirus (BHV-1) gD glycoprotein (GenBank sequence accession No. = L26360) in the form of a 338 bp PCR fragment. After purification, this fragment was digested with PstI and NdeI in order to isolate a 317 bp PstI-NdeI fragment (fragment A).

25 The plasmid pBHV001 (P. Leung-Tack et al., Virology, 1994, 199, 409-421) was digested with NdeI and StyI in order to liberate a 942 bp fragment containing the 3' part of the gene encoding the BHV-1 gD glycoprotein (fragment B).

30 The fragments A and B were ligated together with the vector pVR1012 (Example 6), previously digested with PstI and XbaI, to give the plasmid pAB087 (6134 bp) (Figure No. 4).

Example 9: Construction of the plasmid pAB011 (BRSV F gene)

An RT-PCR reaction according to the technique

described in Example 5 was carried out with the genomic RNA from the bovine respiratory syncytial virus (BRSV) (391-2 strain) (R. Lerch et al., *Virology*, 1991, 181, 118-131), prepared as indicated in Example 3, and 5 with the following oligonucleotides:
AB026 (33 mer) (SEQ ID No. 6)
5' AAAACTGCAGGGATGGCGGCAACAGCCATGAGG 3'
AB027 (31 mer) (SEQ ID No. 7)
5' CGCGGATCCTCATTACTAAAGGAAAGATTG 3'
10 so as to isolate the gene encoding the F fusion glycoprotein (BRSV F) in the form of a 1734 bp PCR fragment. After purification, this fragment was digested with PstI and BamHI in order to isolate a 1717 bp PstI-BamHI fragment. This fragment was ligated with the vector
15 pVR1012 (Example 6), previously digested with PstI and BamHI, to give the plasmid pAB011 (6587 bp) (Figure No. 5).

20 **Example 10: Construction of the plasmid pAB012 (BRSV G gene)**

An RT-PCR reaction according to the technique described in Example 5 was carried out with the genomic RNA from the bovine respiratory syncytial virus (BRSV) (391-2 strain) (R. Lerch et al., *J. Virology*, 1990, 64, 25 5559-5569) and with the following oligonucleotides:
AB028 (32 mer) (SEQ ID No. 8)
5' AAAACTGCAGATGTCCAACCATAACCCATCATC 3'
AB029 (35 mer) (SEQ ID No. 9)
5' CGCGGATCCCTAGATCTGTGTAGTTGATTGATTG 3'
30 so as to isolate the gene encoding the G protein (BRSV G) in the form of a 780 bp PCR fragment. After purification, this fragment was digested with PstI and BamHI in order to isolate a 763 bp PstI-BamHI fragment. This fragment was ligated with the vector pVR1012
35 (Example 6), previously digested with PstI and BamHI, to give the plasmid pAB012 (5634 bp) (Figure No. 6).

Example 11: Construction of the plasmid pAB058 (BVDV C gene)

An RT-PCR reaction according to the technique described in Example 5 was carried out with the genomic RNA from the bovine viral diarrhoea virus (BVDV) (Osloss strain) (L. De Moerlooze et al., J. Gen. Virol., 1993, 74, 1433-1438), prepared according to the technique described in Example 3, and with the following oligonucleotides:

5 AB110 (35 mer) (SEQ ID No. 10)
5' AAAACTGCAGATGTCCGACACAAAAGCAGAAGGGG 3'

10 AB111 (47 mer) (SEQ ID No. 11)
5' CGCGGATCCTCAATAAAATCATTCCACTGCGACTTGAAACAAAAC 3'

15 so as to amplify a 342 bp fragment containing the gene encoding the C capsid protein from the BVDV virus. After purification, the RT-PCR product was digested with PstI and BamHI to give a 324 bp PstI-BamHI fragment.

This fragment was ligated with the vector pVR1012 (Example 6), previously digested with PstI and BamHI, to give the plasmid pAB058 (5183 bp) (Figure 20 No. 7).

Example 12: Construction of the plasmid pAB059 (BVDV E1 "gene")

An RT-PCR reaction according to the technique described in Example 5 was carried out with the genomic RNA from the bovine viral diarrhoea virus (BVDV) (Osloss strain) (L. De Moerlooze et al., J. Gen. Virol., 1993, 74, 1433-1438) and with the following oligonucleotides:

25 AB114 (32 mer) (SEQ ID No. 12)
5' ACGCGTCGACATGAAGAAACTAGAGAAAGCCC 3'
AB115 (33 mer) (SEQ ID No. 13)
5' CGCGGATCCTCAGCCGGTTGCAAACGGGAG 3'
so as to isolate the sequence encoding the BVDV virus
35 E1 protein in the form of a 1381 bp PCR fragment. After purification, this fragment was digested with SalI and BamHI to give a 1367 bp SalI-BamHI fragment.

This fragment was ligated with the vector pVR1012 (Example 6), previously digested with SalI and

BamHI, to give the plasmid pAB059 (6236 bp) (Figure No. 8).

Example 13: Construction of the plasmid pAB060 (BVDV E2 "gene")

5 An RT-PCR reaction according to the technique described in Example 5 was carried out with the genomic RNA from the bovine viral diarrhoea virus (BVDV) (Osloss strain) (L. De Moerlooze et al., J. Gen. Virol., 1993, 74, 1433-1438) and with the following 10 oligonucleotides:

AB116 (36 mer) (SEQ ID No. 14)

5' ACGCGTCGACATGACGACTACTGCATTCCCTGGTATG 3'

AB117 (33 mer) (SEQ ID No. 15)

5' CGCGGATCCTCATTGACGTCCCGAGGTCATTTG 3'

15 so as to isolate the sequence encoding the BVDV virus E2 protein in the form of a 1252 bp PCR fragment. After purification, this fragment was digested with SalI and BamHI to give a 1238 bp SalI-BamHI fragment.

20 This fragment was ligated with the vector pVR1012 (Example 6), previously digested with SalI and BamHI, to give the plasmid pAB060 (6107 bp) (Figure No. 9).

25 **Example 14: Construction of the plasmid pAB071 (BPIV HN gene)**

An RT-PCR reaction according to the technique described in Example 5 was carried out with the genomic RNA from the type 3 bovine parainfluenza virus (PI3 = BPIV) and with the following oligonucleotides:

30 AB130 (33 mer) (SEQ ID No. 16)

5' TTTGTCGACATGGAATATTGGAAACACACACAAAC 3'

AB131 (33 mer) (SEQ ID No. 17)

5' TTTGGATCCTTAGCTGCAGTTTCGGAACCTTC 3'

35 so as to isolate the gene encoding the BPIV HN glycoprotein (HN gene sequence deposited by H. Shibuta in 1987. GenBank sequence accession No. = Y00115) in the form of a 1737 bp PCR fragment. After purification, this fragment was digested with SalI and BamHI in order to isolate a 1725 bp SalI-BamHI fragment. This fragment

was ligated with the vector pVR1012 (Example 6), previously digested with SalI and BamHI, to give the plasmid pAB071 (6593 bp) (Figure No. 10).

5 **Example 15: Construction of the plasmid pAB072 (BPIV F gene)**

An RT-PCR reaction according to the technique described in Example 5 was carried out with the genomic RNA from the type 3 bovine parainfluenza virus (PI3 = 10 BPIV) and with the following oligonucleotides:

AB132 (30 mer) (SEQ ID No. 18)

5' TTTGTCGACATGATCATCACAAACACAATC 3'

AB133 (30 mer) (SEQ ID No. 19)

5' TTTGGATCCTCATTGTCTACTTGTAGTAC 3'

15 so as to isolate the gene encoding the BPIV F glycoprotein (F gene sequence deposited by H. Shibuta in 1987. GenBank sequence accession No. = Y00115) in the form of a 1641 bp PCR fragment. After purification, this fragment was digested with SalI and BamHI in order 20 to isolate a 1629 bp SalI-BamHI fragment. This fragment was ligated with the vector pVR1012 (Example 6), previously digested with SalI and BamHI, to give the plasmid pAB072 (6497 bp) (Figure No. 11).

25 **Example 16: Preparation and purification of the plasmids**

For the preparation of the plasmids intended for the vaccination of animals, any technique may be used which makes it possible to obtain a suspension of 30 purified plasmids predominantly in a supercoiled form. These techniques are well known to persons skilled in the art. There may be mentioned in particular the alkaline lysis technique followed by two successive ultracentrifugations on a caesium chloride gradient in 35 the presence of ethidium bromide as described in J. Sambrook et al. (*Molecular Cloning: A Laboratory Manual*, 2nd edition, Cold Spring Harbor Laboratory, Cold Spring Harbor, New York, 1989). Reference may also be made to Patent Applications PCT WO 95/21250 and PCT WO

96/02658, which describe methods for producing, on an industrial scale, plasmids which can be used for vaccination. For the purposes of the manufacture of vaccines (see Example 17), the purified plasmids are 5 resuspended so as to obtain solutions at a high concentration (> 2 mg/ml) which are compatible with storage. To do this the plasmids are resuspended either in ultrapure water or in TE buffer (10 mM Tris-HCl; 1 mM EDTA, pH 8.0).

10

Example 17: Manufacture of the associated vaccines

The various plasmids necessary for the manufacture of an associated vaccine are mixed starting with their concentrated solutions (Example 16). The 15 mixtures are prepared such that the final concentration of each plasmid corresponds to the effective dose of each plasmid. The solutions which can be used to adjust the final concentration of the vaccine may be either a 0.9% NaCl solution, or PBS buffer.

20

Specific formulations such as liposomes or cationic lipids, may also be used for the manufacture of the vaccines.

Example 18: Vaccination of bovines

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The bovines are vaccinated with doses of 100 µg, 250 µg or 500 µg per plasmid. The injections are performed with a needle by the intramuscular route either at the level of the gluteus muscle, or at the level of the neck muscles. The vaccinal doses are 30 administered in volumes of between 1 and 5 ml.

CLAIMS

1. Bovine vaccine formula against bovine respiratory pathology, comprising at least three polynucleotide vaccine valencies each comprising a plasmid integrating, so as to express it in vivo in the host cells, a gene with one bovine respiratory pathogen valency, these valencies being selected from the group consisting of bovine herpesvirus, bovine respiratory syncytial virus, mucosal disease virus and type 3 parainfluenza virus, the plasmids comprising, for each valency, one or more of the genes selected from the group consisting of gB and gD for the bovine herpesvirus, F and G for the bovine respiratory syncytial virus, E2, C + E1 + E2 and E1 + E2 for the mucosal disease virus, HN and F for the type 3 parainfluenza virus.
2. Vaccine formula according to Claim 1, which comprises the four polynucleotide vaccine valencies.
3. Vaccine formula according to Claim 1 or 2, which comprises the bovine herpesvirus gB and gD genes, in the same plasmid or in different plasmids.
4. Formula according to Claim 1 or 2, which comprises the bovine respiratory syncytial virus F and G genes, in the same plasmid or in different plasmids.
5. Vaccine formula according to Claim 1 or 2, wherein the plasmid for the mucosal disease virus comprises the E2 gene.
6. Vaccine formula according to Claim 1 or 2, wherein, for the type 3 parainfluenza virus valency, it comprises the HN gene in one plasmid or all the genes encoding HN and F in the same plasmid or in different plasmids.
7. Vaccine formula according to all of Claims 1 to 6.
8. Vaccine formula according to any one of Claims 1 to 7, which comprises from 10 ng to 1 mg, preferably from 100 ng to 500 μ g, still more preferably from 1 μ g to 250 μ g of each plasmid.

9. Use of one or more plasmids as described in any one of Claims 1 to 8, for the manufacture of a vaccine intended to vaccinate bovines first vaccinated by means of a first vaccine selected from the group consisting of a live whole vaccine, an inactivated whole vaccine, a subunit vaccine, a recombinant vaccine, this first vaccine having the antigen(s) encoded by the plasmid(s) or antigen(s) providing cross-protection.

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10. Vaccination kit grouping together a vaccine formula according to any one of Claims 1 to 8, and a vaccine selected from the group consisting of a live whole vaccine, an inactivated whole vaccine, a subunit vaccine, a recombinant vaccine, this first vaccine having the antigen encoded by the polynucleotide vaccine or an antigen providing cross-protection, for an administration of the latter in first vaccination and as a booster with the vaccine formula.

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20. Vaccine formula according to any one of Claims 1 to 8, accompanied by a leaflet indicating that this formula can be used as a booster for a first vaccine selected from the group consisting of a live whole vaccine, an inactivated whole vaccine, a subunit vaccine, a recombinant vaccine, this first vaccine having the antigen encoded by the polynucleotide vaccine or an antigen providing cross-protection.

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COMPANY CALLED: RHONE MERIEUX

POLYNUCLEOTIDE VACCINE FORMULA IN PARTICULAR AGAINST
BOVINE RESPIRATORY PATHOLOGY

ABSTRACT OF THE CONTENT OF THE INVENTION

The bovine vaccine formula against bovine respiratory pathology comprises at least three polynucleotide vaccine valencies each comprising a plasmid integrating, so as to express it in vivo in the host cells, a gene with one bovine respiratory pathogen valency, these valencies being selected from those of the group consisting of bovine herpesvirus, bovine respiratory syncytial virus, mucosal disease virus and type 3 parainfluenza virus, the plasmids comprising, for each valency, one or more of the genes selected from the group consisting of gB and gD for the bovine herpesvirus, F and G for the bovine respiratory syncytial virus, E2, C + E1 + E2 and E1 + E2 for the mucosal disease virus, and HN and F for the type 3 parainfluenza virus.

Figure 4.



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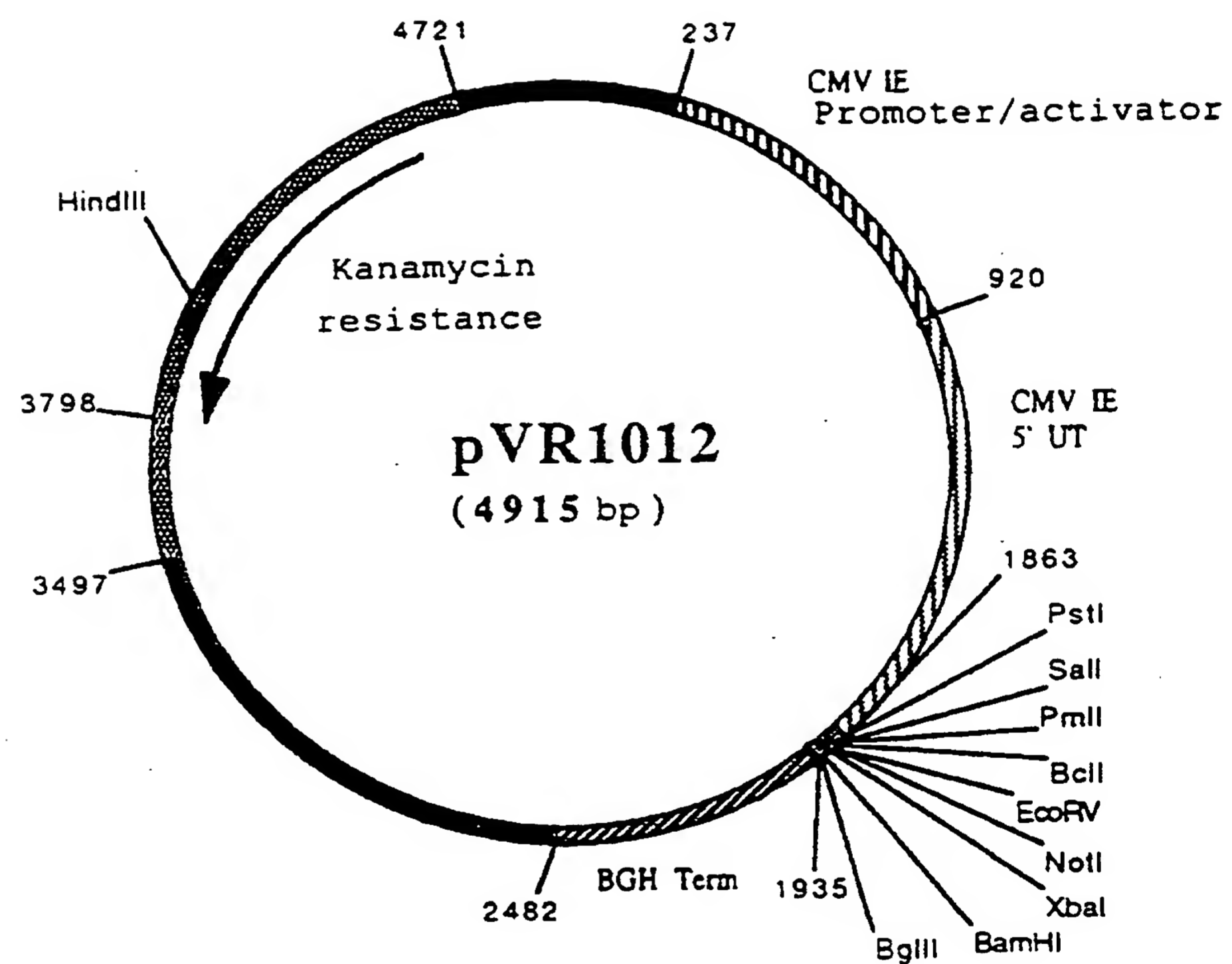


Figure No. 1



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1 ATGGCCGCTCGCGGCGGTGCTGAACGGCGCCGGCGCCGGAGACGGTGGCGAGGACAGCGT
1 Met Ala Ala Arg Gly Gly Ala Glu Arg Ala Ala Gly Ala Gly Asp Gly Arg Arg Gly Gln Arg

64 CGTCATCTACGACCGGGACGTGTTCTGCTCTACGGCTCTGCAGCGCCTGGCGCCGGC
22 Arg His Leu Arg Pro Gly Arg Val Leu Ala Ala Leu Arg Gly Pro Ala Ala Pro Gly Ala Gly

127 GGGCGCCGCCGCGCTAGCCGCTGCCCTGCTATGGCGACGTGGGCCCTGCTGCTGGCGCCG
43 Gly Ala Arg Ala Ala Leu Ala Ala Leu Leu Trp Ala Thr Trp Ala Leu Leu Leu Ala Ala

190 CCCGCCGCGGGGGACCGGGGACAACGCCCGGGCCCCCGCCGAAGAGGCCGCGAGCCCG
64 Pro Ala Ala Gly Arg Pro Ala Thr Thr Pro Pro Ala Pro Pro Glu Glu Ala Ala Ser Pro

253 GCGCCCCCGCGAGCCCCAGCCCCCGGGCCCCGACGGCGACGACGCCGCCAGCCCCGACAAC
85 Ala Pro Pro Ala Ser Pro Ser Pro Pro Gly Pro Asp Gly Asp Asp Ala Ala Ser Pro Asp Asp

316 AGCACAGACGTGGCGCCGCCGCTCCGGCTCGCGAGCCGGGGAAAAACTCGCGCTTC
106 Ser Thr Asp Val Arg Ala Ala Leu Arg Leu Ala Gln Ala Ala Gly Glu Asn Ser Arg Phe Phe

379 GTGTGCCCGCCGCCCTCGGGCGCCACGGTGGTCCGGCTCGCGCCCGGCCGTGCCCTGAC
127 Val Cys Pro Pro Pro Ser Gly Ala Thr Val Val Arg Leu Ala Pro Ala Arg Pro Cys Pro Glu

442 TACGGGCTCGGGCGGA ACTACACGGAGGGCATCGCGTCATTACAAGGAGAACATCGCGCCG
148 Tyr Gly Leu Gly Arg Asn Tyr Thr Glu Gly Ile Gly Val Ile Tyr Ile Glu Asn Ile Ala Pro

505 TACACGTTCAAGGCCTACATTACAAAAACGTGATCGTGACCACGACCTGGCGGGCAGCACG
169 Tyr Thr Phe Lys Ala Tyr Ile Tyr Lys Asn Val Ile Val Thr Thr Trp Ala Gly Ser Thr

568 TACGGCGGCCATTACAAACCAGTACACGGACCGCGTGGCGATGGCGAGATCACCGAC
190 Tyr Ala Ala Ile Thr Asn Gln Tyr Thr Asp Arg Val Pro Val Gly Met Gly Glu Ile Thr Asp

631 CTGGTGGACAAGAAGTGGCGCTGCCCTTCGAAAGCCGAGTACCTGGCGAGCGGGCGCAAGGTC
211 Leu Val Asp Lys Lys Trp Arg Cys Leu Ser Ile Ala Glu Tyr Leu Arg Ser Gly Arg Lys Val

694 GTGGCCTTGACCGCGACGACGACCCCTGGGAGGCCGCTGAAGCCTCCGCGCTGAGCGCG
232 Val Ala Phe Asp Arg Asp Asp Pro Trp Glu Ala Pro Leu Lys Pro Ala Arg Leu Ser Ala

757 CCCGGGGTGCAGGGCTGGCACACGACGGACGATGTGTACACGGCGCTGGCTGGCGGGCTC
253 Pro Gly Val Arg Gly Trp His Thr Thr Asp Val Tyr Thr Ala Leu Gly Ser Ala Gly Leu

820 TACCGCACGGGCACCTCTGTGAACCGCATCGTGGAAAGAAGTGGAGGGCGCTCGGTGTACCCG
274 Tyr Arg Thr Gly Thr Ser Val Asn Cys Ile Val Glu Glu Val Ala Arg Ser Val Tyr Pro

883 TACGACTCGTTGGCGCTCTGACCCGGGACATTATCTACATGTCGCCCTTACGGCGTGC
295 Tyr Asp Ser Phe Ala Leu Ser Thr Gly Asp Ile Ile Tyr Met Ser Pro Phe Tyr Gly Leu Arg

946 GAGGGCGCCACCGCGAGCACACCAGGCTACTGCCGGAGCGCTTCAGCAGATCGAGGGCTA
316 Glu Gly Ala His Arg Glu His Thr Arg Leu Leu Ala Gly Ala Leu Pro Ala Asp Arg Gly Leu

1009 CTACAAGCGCGACATGCCACGGCGCCCTCAAGGAGCCGGTCTCGCCGAACCTTTGCG
337 Leu Gln Ala Arg His Gly His Gly Pro Ala Pro Gln Gly Ala Gly Leu Ala Glu Leu Phe Ala

1072 TACACAGCACGTGACGGTAGCCTGGACTGGTCCCCAAGCCCACCGTGTGCTCCCTGGC
358 Tyr Thr Ala Arg Asp Gly Ser Leu Gly Leu Gly Ala Gln Ala Gln Lys Arg Val Leu Ala Gly



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1135 CAAGTGGCGCGAGGCGGACGAAATGCTGGAGACGAGAGCCGCGGGAACTTCCGCTTCACGGC
379► GlnValAlaArgGlyGlyArgAsnAlaAlaArgArgGluProArgGluLeuProLeuHisGly
1198 CCGCTCGCTCTCGCGACCTTGTGAGCGACAGCCACACCTTCGCGTTGCAGAATGTGCCGCT
400► ProLeuAlaLeuGlyAspLeuCysGluArgGlnProHisLeuArgValAlaGluCysAlaAla
1261 GAGCGACTGCGTGATCGAAGAGGCCGAGGCCGCGTCGAGCCGCTACCGCGAGCCTACAA
421► GluArgLeuArgAspArgArgGlyArgGlyArgGlyArgAlaArgLeuProArgAlaLeuGln
1324 CGGCACGCCACGTGCTGTCGGCAGCTGGAGACGTACCTGGCGCCGGCGCTTGTGCGTGGC
442► ArgHisAlaArgAlaValGlyGlnLeuGlyAspValProGlyAlaArgArgLeuCysArgGly
1387 CTTCCGGCGATGCTCAGCAACGAGCTGGCCAAGCTGTACCTGCAGGAGCTGGCGCGCTCGAAC
463► LeuProAlaMetLeuSerAsnGluLeuAlaLysLeuTyrLeuGlnGluLeuAlaArgSerAsn
1450 CGCACGCTCGAGGGCTTTCGCCGCCGCCGCCAAGCCGGCCGGCGCGCGCGCGCGCGCG
484► GlyThrLeuGluGlyLeuPheAlaAlaAlaAlaProLysProGlyProArgArgAlaArgArg
1513 GCCGCGCCGTCTGCCGCCGCCGCCGCCGCCAACGGGCCCGCCGACGGCGACGCC
505► AlaAlaProSerAlaProGlyGlyProGlyAlaAlaAsnGlyProAlaGlyAspGlyAspAla
1576 GCGGGCGCGGTGACTACCGTGAGCTGGCCAGTTGCCGGCTGCAGTTACCTACGACCAC
526► GlyGlyArgValThrThrValSerSerAlaGluPheAlaAlaLeuGlnPheThrTyrAspHis
1639 ATCCAGGACCACGTGAACACCATGTTAGCCGCTGGCACGTCCTGGTGCCTGCTGCAGAAC
547► IleGlnAspHisValAsnThrMetPheSerArgLeuAlaThrSerTrpCysLeuLeuGlnAsn
1702 AAGGAGCGCCGCCCTGTGGCCGAGGCGGCTAACGCTAACCCAGCGCGGCCAGCGCTGG
568► LysGluArgAlaLeuTrpAlaGluAlaAlaLysLeuAsnProSerAlaAlaAlaSerAlaAla
1765 CTGGACCGCCGCCGCCTGGCCGAGGCGGCTAACGCTAACCCAGCGCGGCCAGCGCTGG
589► LeuAspArgArgAlaAlaAlaArgMetLeuGlyAspAlaMetAlaValThrTyrCysHisGlu
1828 CTGGCGAGGGCGCGTGTTCATCGAGAACTCGATGCCGCCGCCGGCGTTGCTACAGC
610► LeuGlyGluGlyArgValPheIleGluAsnSerMetArgAlaProGlyGlyValCysTyrSer
1891 CGCCCGCCGGCTCCCTTGCCTTCGGCAACGAGAGCGAGCCGGTGGAGGGCCAGCTCGCGAG
631► ArgProProValSerPheAlaPheGlyAsnGluSerGluProValGluGlyGlnLeuGlyGlu
1954 GACAACGAGCTGCCGCCGCCGCCGAGCTCGGAGCCCTGCACGCCAACACACAGCGCTAC
652► AspAsnGluLeuLeuProGlyArgGluLeuValGluProCysThrAlaAsnHisLysArgTyr
2017 TTCCGCTTGGCCGGACTACGTGACTACGAGAACTACGGTACGTGGCGGGTCCGCTC
673► PheArgPheGlyAlaAspTyrValTyrTyrGluAsnTyrAlaTyrValArgArgValProLeu
2080 GCGGAGCTGGAGGTGATCAGCACCTTGTGGACCTAACCTCACGGTTCTGGAGGACCGCGAG
694► AlaGluLeuGluValIleSerThrPheValAspLeuAsnLeuThrValLeuGluAspArgGlu
2143 TTCTGCCGCTAGAAAGTGTACACGCCGCCGAGCTCGCCGACACGGGTCTGCTGACTACAGC
715► PheLeuProLeuGluValTyrThrArgAlaGluLeuAlaAspThrGlyLeuLeuAspTyrSer
2206 GAGATACAGCCCCGCAACCAGCTGCACGAGCTCCGGTCTACGACATTGACCCGTGGTCAAG
736► GluIleGlnArgArgAsnGlnLeuHisGluLeuArgPheTyrAspIleAspArgValValLys



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2269 ACGGACGGCAATATGCCATCGGAGGGCTGCCAACTCTTCAGGGCCTGGGCGCCGTC
757 ▶ Thr Asp Gly Asn Met Ala Ile Met Arg Gly Leu Ala Asn Phe Phe Gln Gly Leu Gly Ala Val
2332 GGGCAGGCGGTGGGCACGGTGGCTGGCGCCGGTGGCGCTCTGACCGTGTGGC
778 ▶ Gly Gln Ala Val Gly Thr Val Val Leu Gly Ala Ala Gly Ala Ala Leu Ser Thr Val Ser Gly
2395 ATGCCCTCGTTATTGCGAACCGTTGGCGCTGGCCACGGGCTGCTGGTGTGCGCCGG
799 ▶ Ile Ala Ser Phe Ile Ala Asn Pro Phe Gly Ala Leu Ala Thr Gly Leu Leu Val Leu Ala Gly
2458 CTGGTGGCCGCTTCTGGCGTACCGGTACATTCCCGCTCCGCAGCAACCCATGAAGGCG
820 ▶ Leu Val Ala Ala Phe Leu Ala Tyr Arg Tyr Ile Ser Arg Leu Arg Ser Asn Pro Met Lys Ala
2521 CTGTACCCGATCACCAACGCCGGCTCAAGGACGACGCCGGCGCAACCGCCCCGGCGAG
841 ▶ Leu Tyr Pro Ile Thr Thr Arg Ala Leu Lys Asp Asp Ala Arg Gly Ala Thr Ala Pro Gly Glu
2584 GAAGAGGAGGAGTTTGACGCCAAACTGGAGCAGGCCGGAGATGATCAAATATGTCG
862 ▶ Glu Glu Glu Glu Phe Asp Ala Ala Lys Leu Glu Gln Ala Arg Glu Met Ile Lys Tyr Met Ser
2647 CTCGTGTCAGCGGTCAGCGGCAAGAGCACAGGCACAAAGGCACAAAGGCAGCCGCTG
883 ▶ Leu Val Ser Ala Val Glu Arg Gln Glu His Lys Ala Lys Lys Ser Asn Lys Gly Gly Pro Leu
2710 CTGGCGACCCGGCTGACGCAGCTGGCGCTCGCGGGAGCGCCGGAGTACCAAGCAGCTT
904 ▶ Leu Ala Thr Arg Leu Thr Gln Leu Ala Leu Arg Arg Arg Ala Pro Pro Glu Tyr Gln Gln Leu
2773 CCGATGCCGACGTCGGGGGGCATGA
925 ▶ Pro Met Ala Asp Val Gly Ala ...

Figure No. 2 (end)



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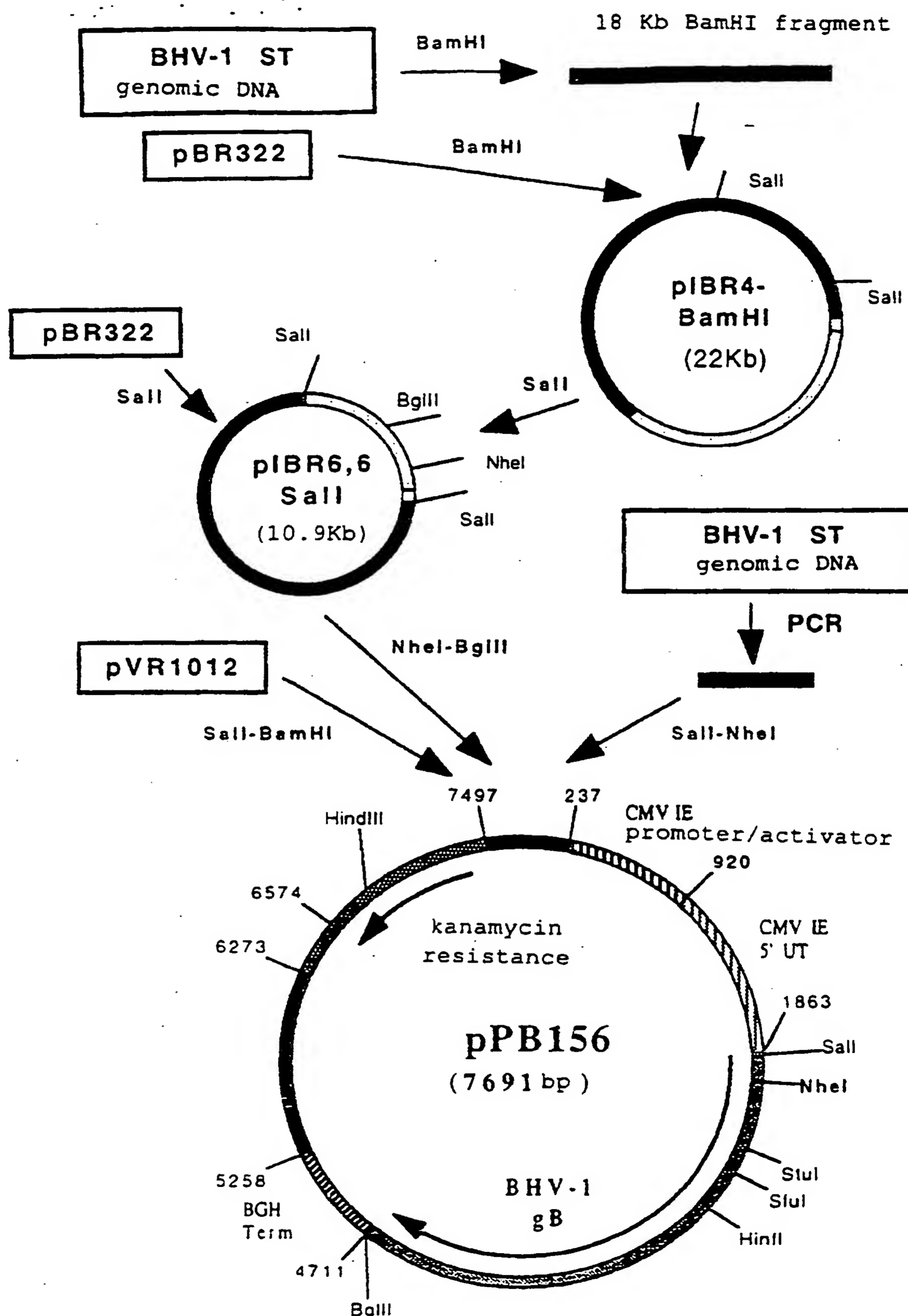
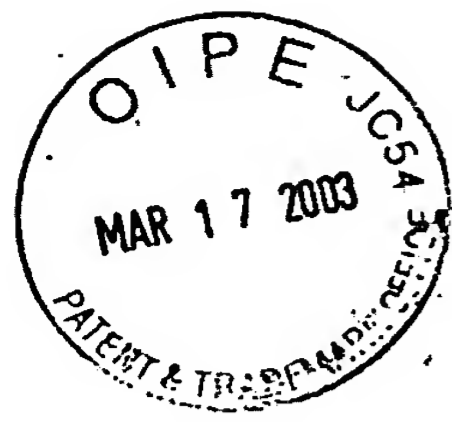


Figure No. 3



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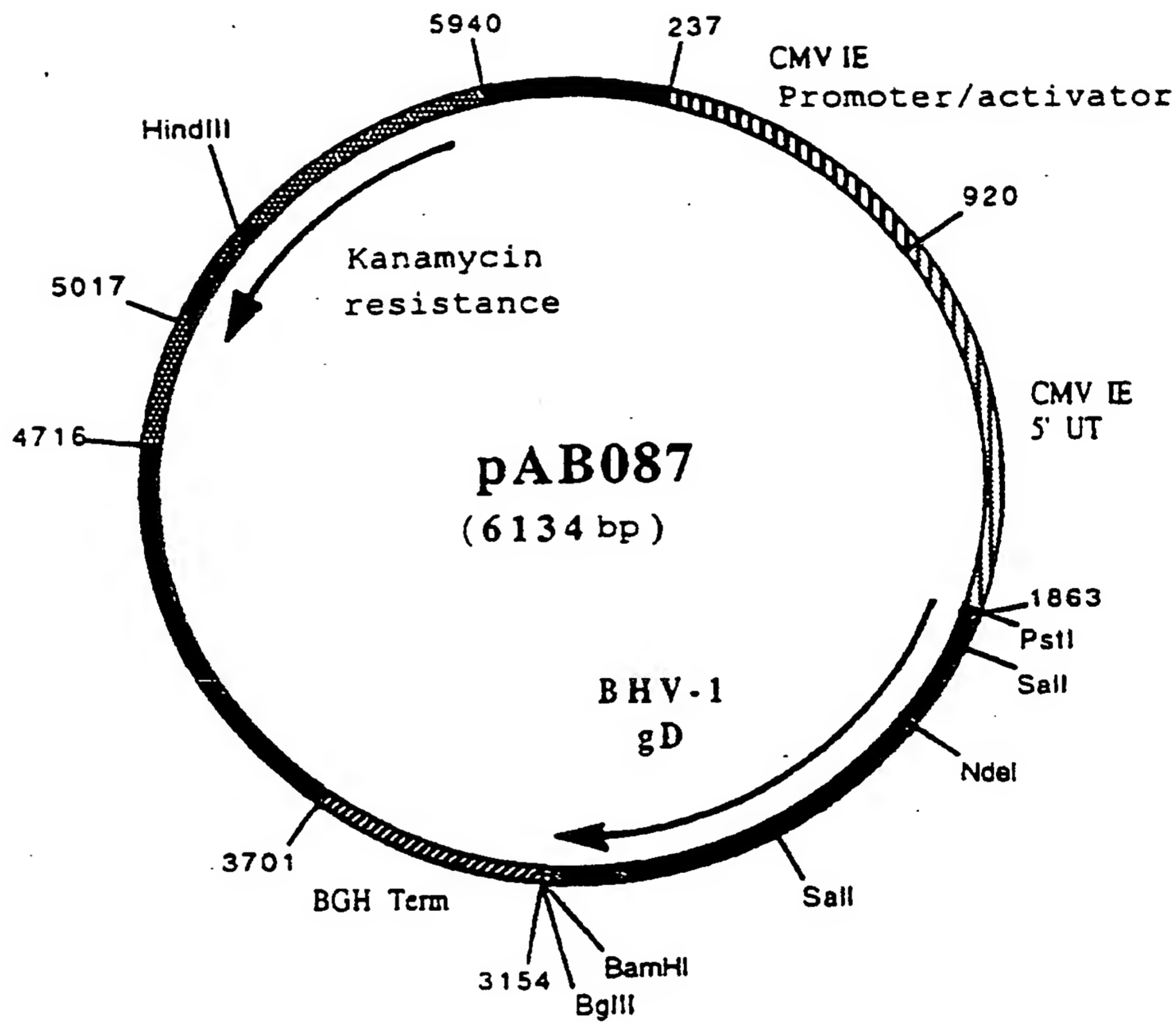


Figure No. 4

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MAR 17 2003
U.S. GOVERNMENT PRINTING OFFICE: 2003 50-2003-000-00000

7113

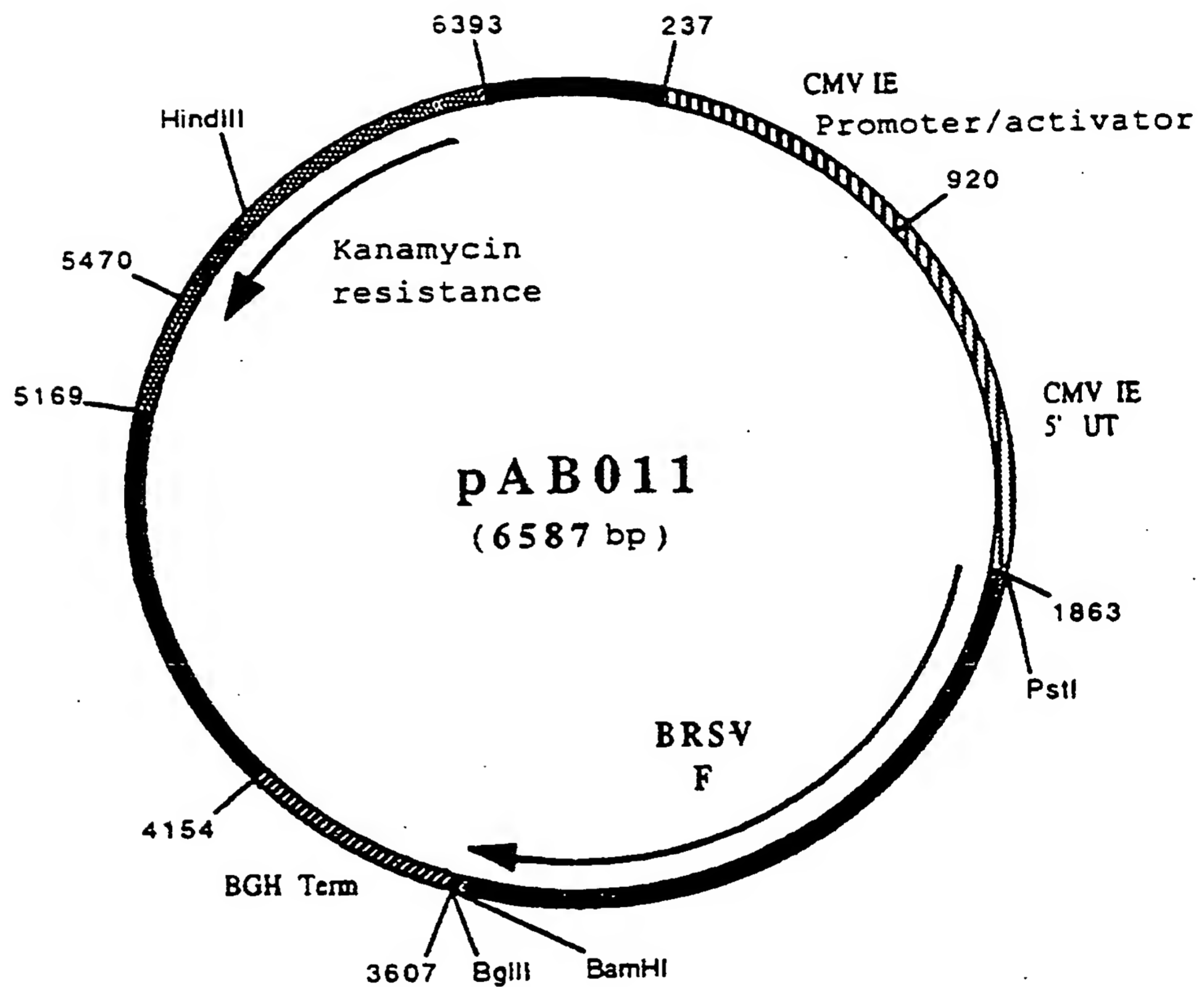


Figure No. 5



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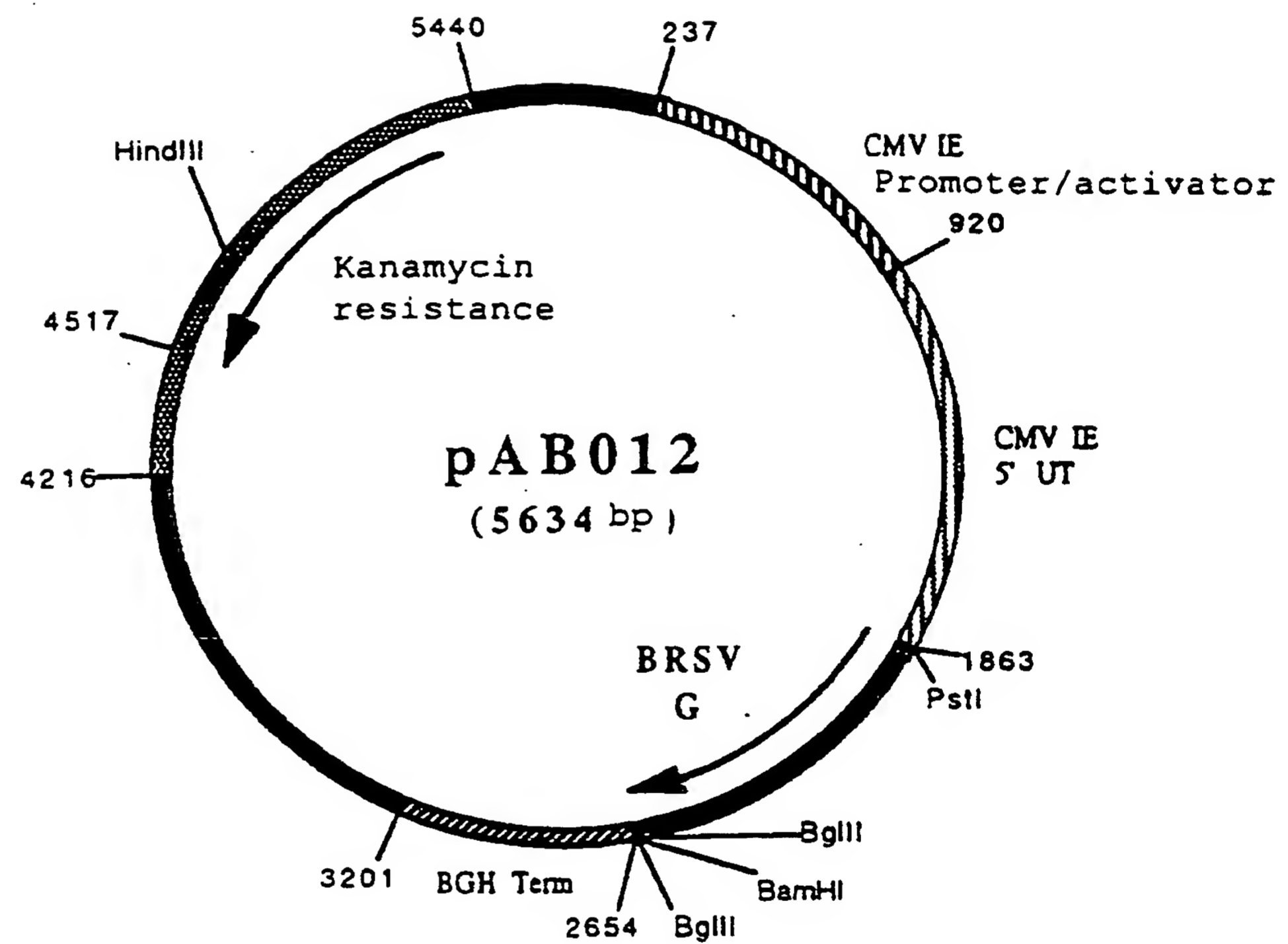


Figure No. 6



913

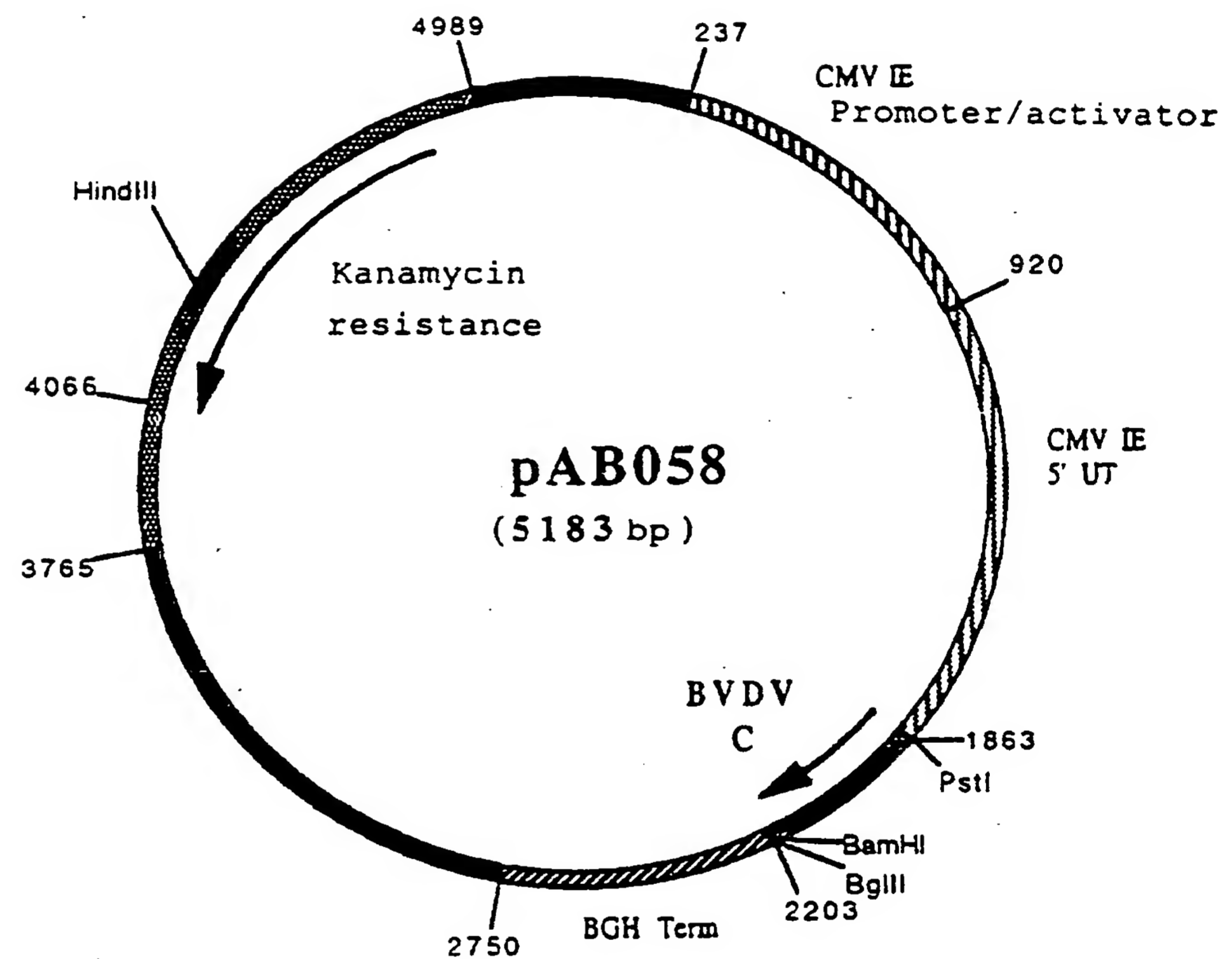


Figure No. 7

O I P E MAR 17 2003 JC54
PRACTICE TRANSMISSIONS

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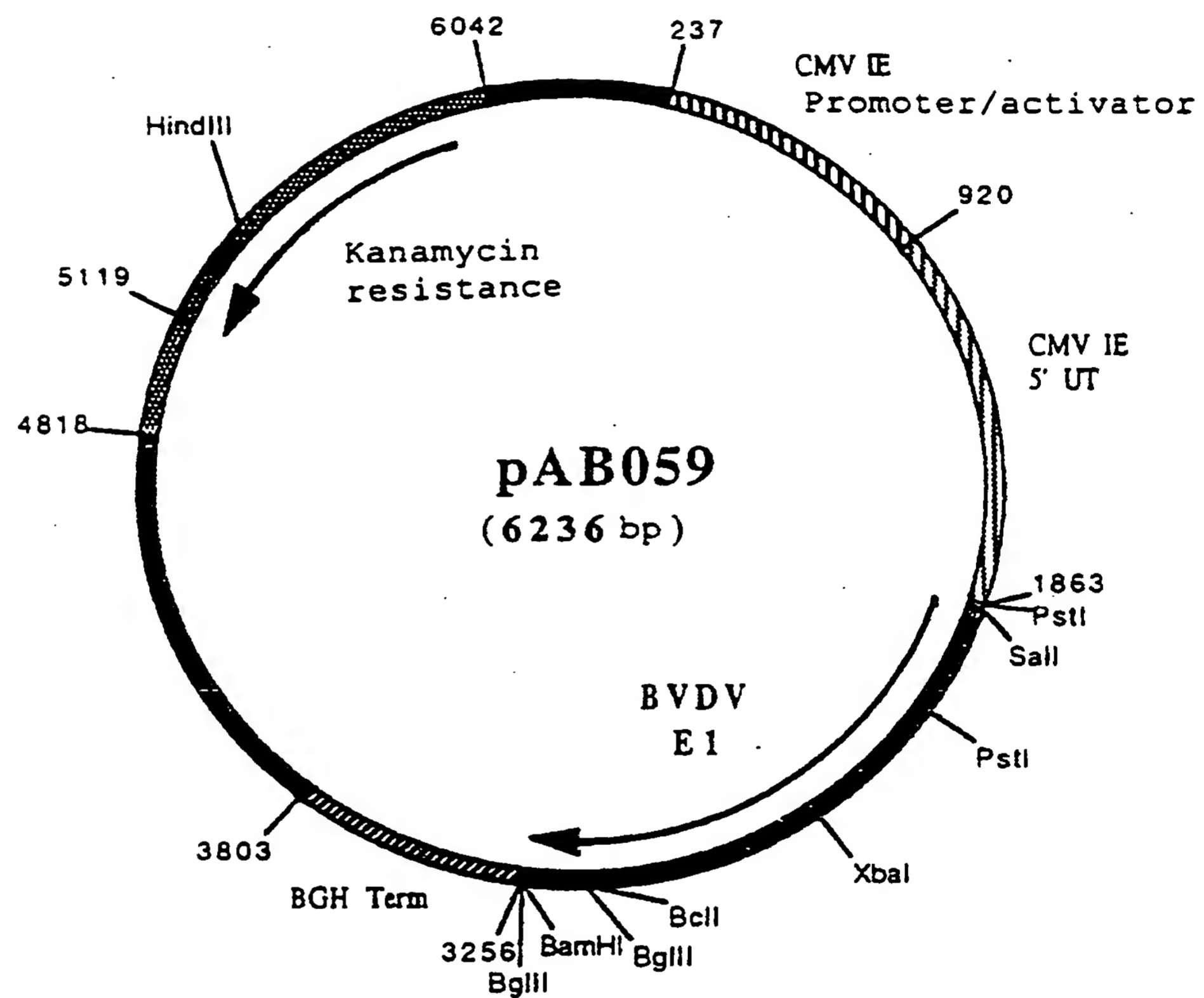


Figure No. 8



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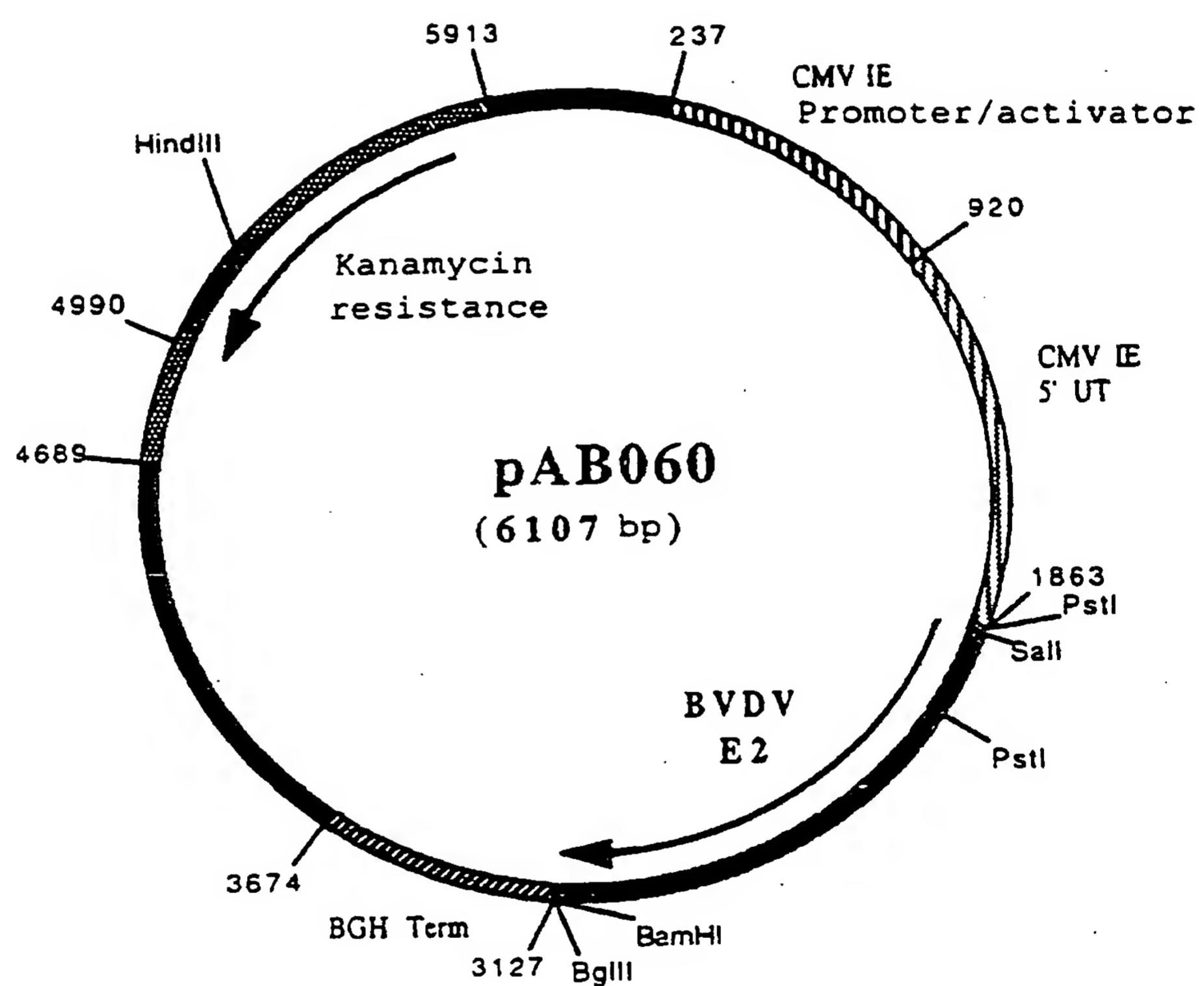


Figure No. 9



12113

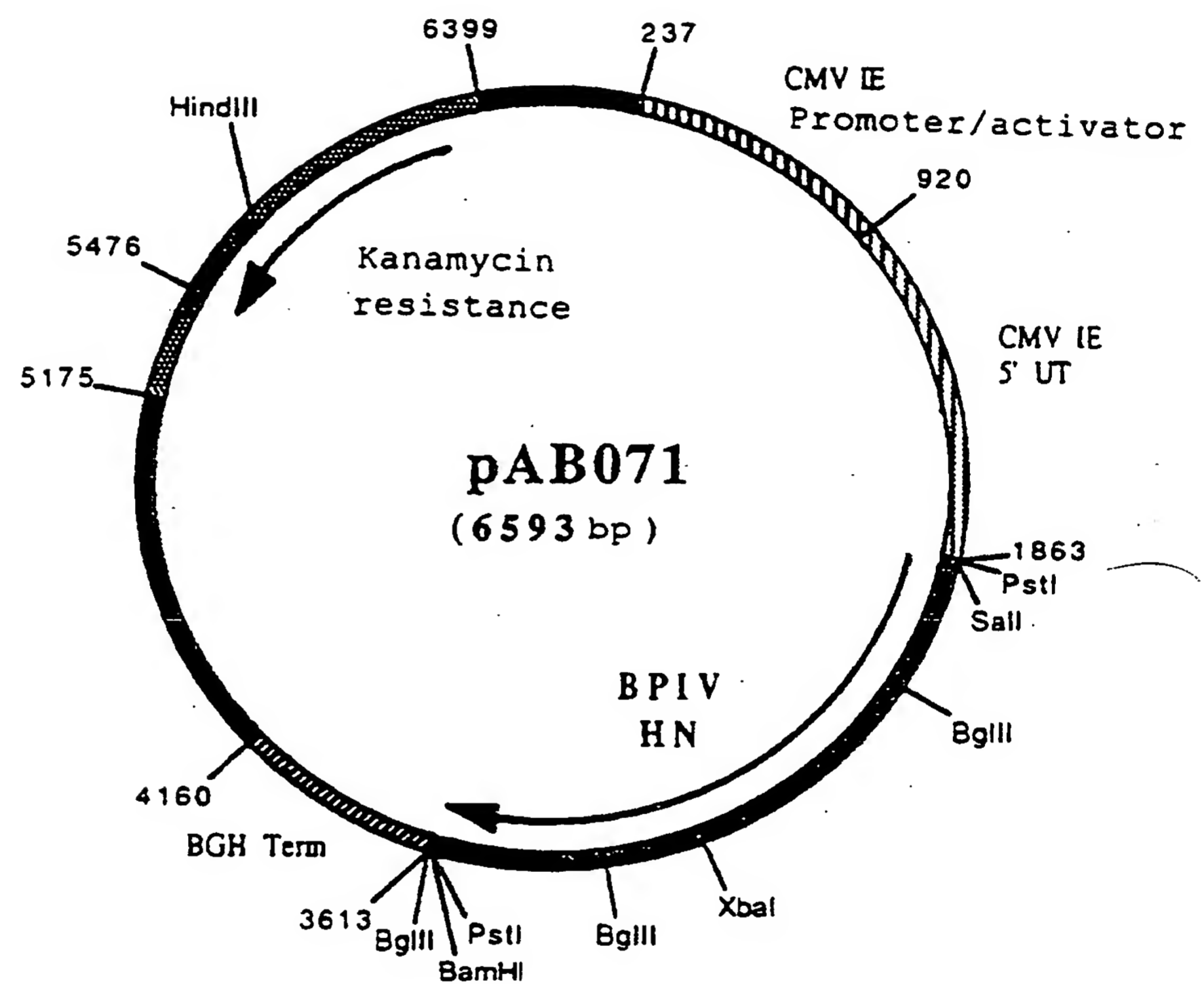


Figure No. 10

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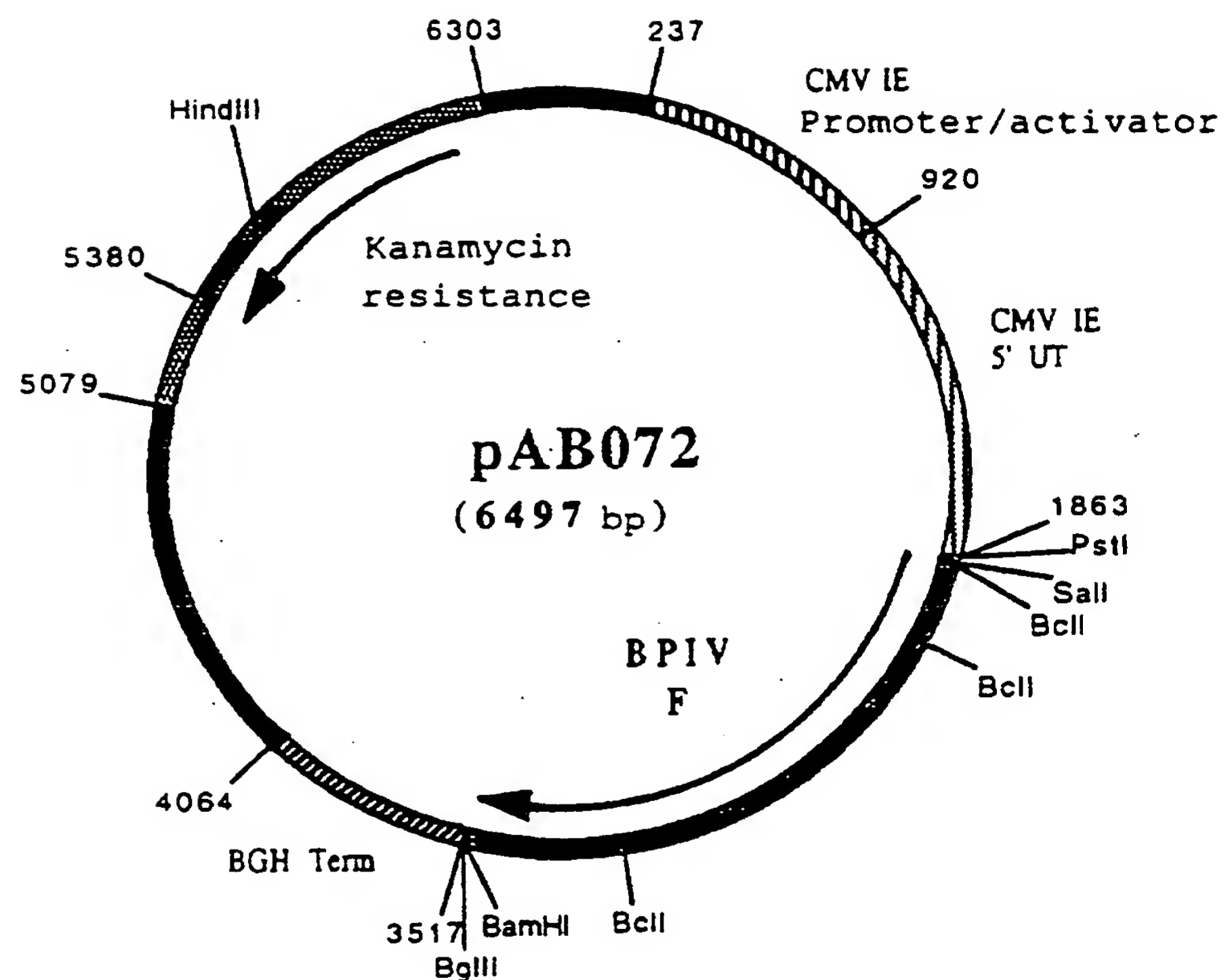


Figure No. 11